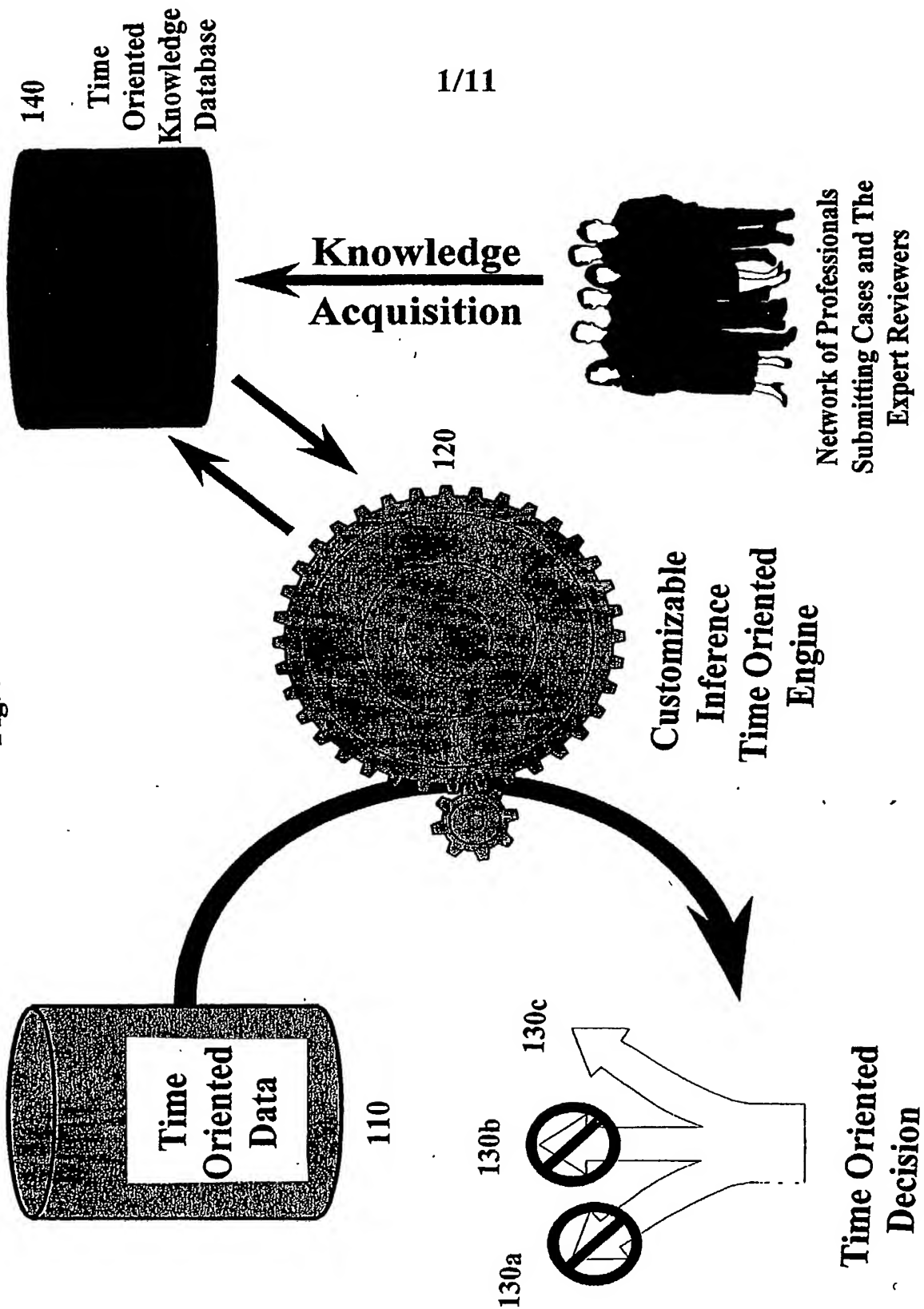
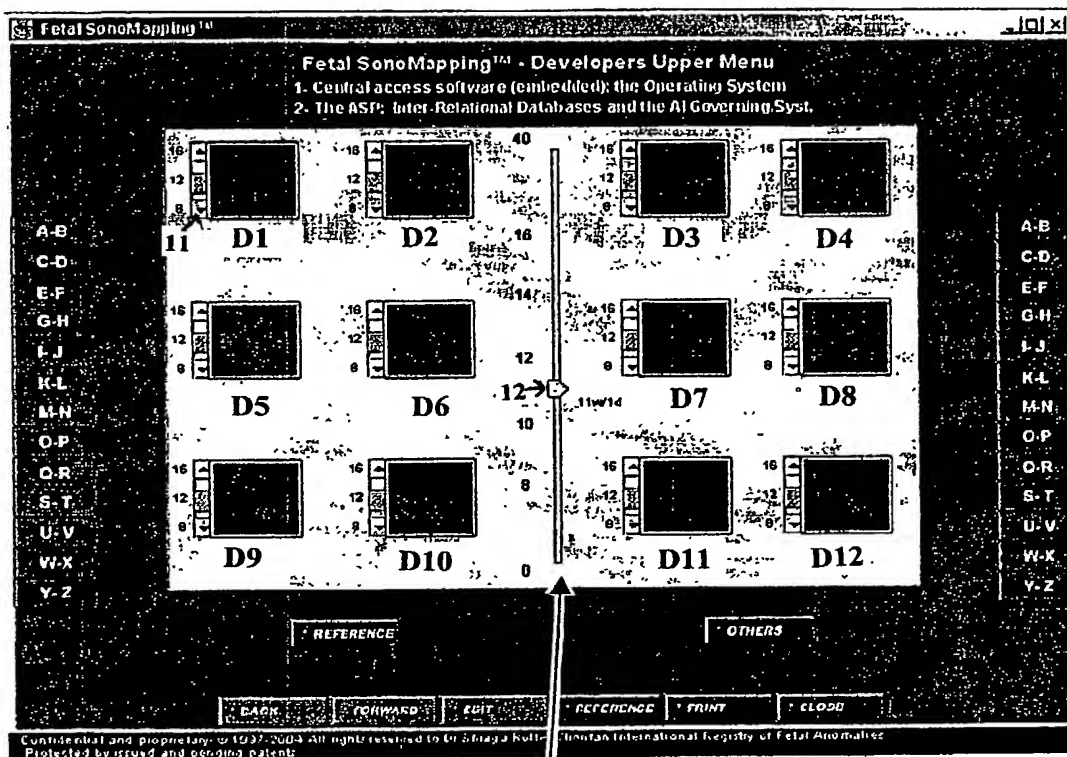


Fig. 1



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Fig. 2

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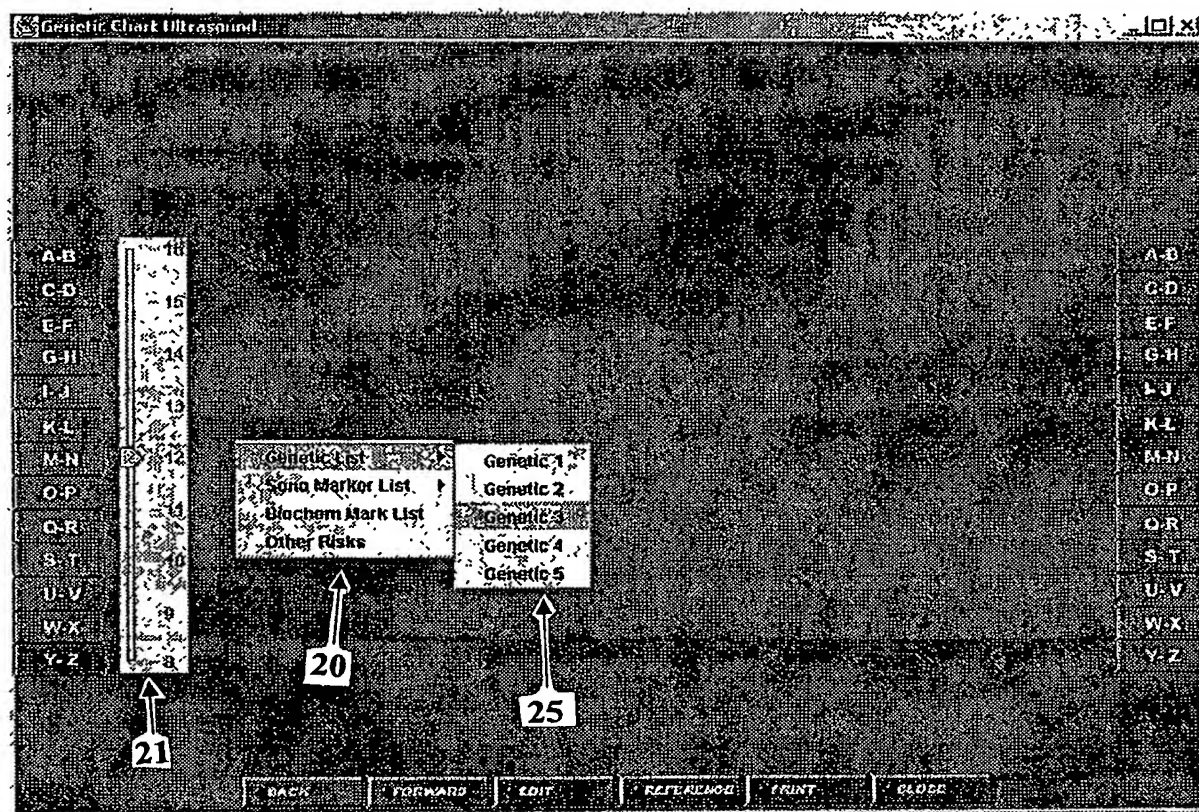


Fig. 2A

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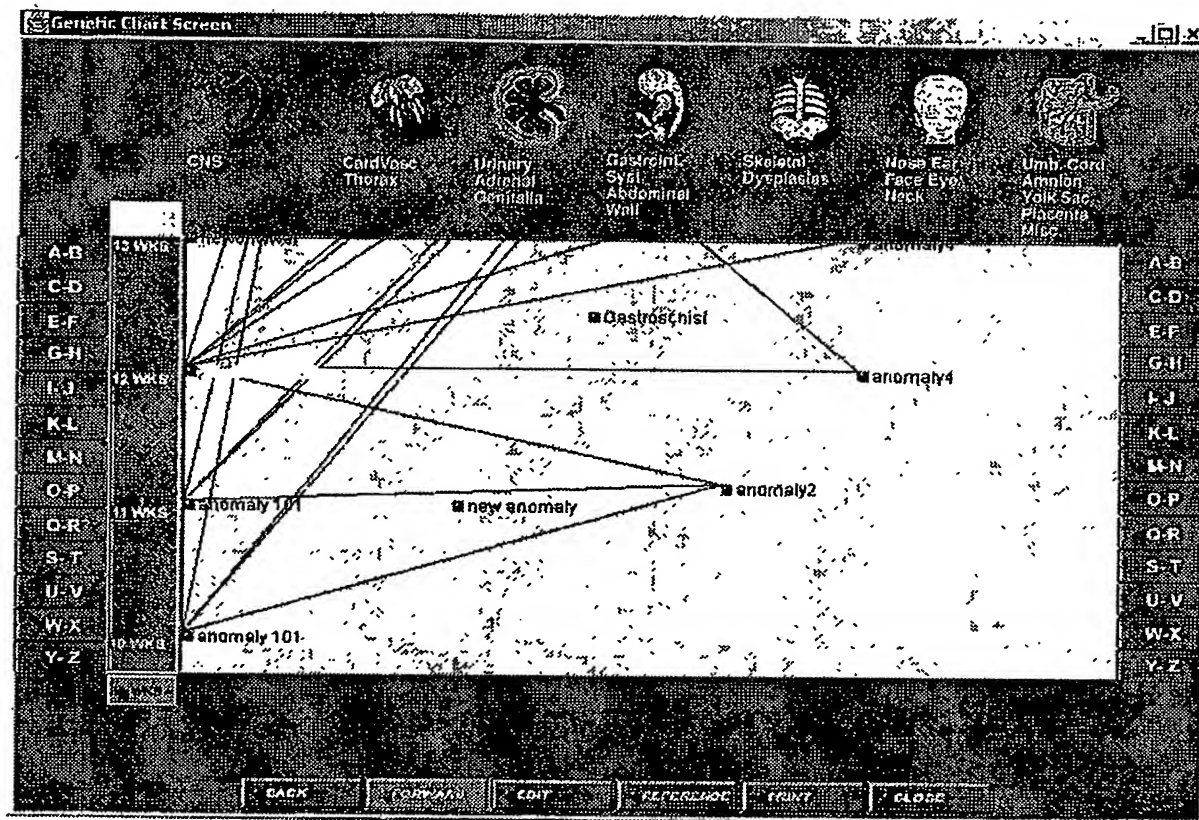


Fig. 2B

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Algo based new associated sign

IRONFAN Anomalies	Earliest seen	Algo Based Most Common Syndrome	Algo Based Associated Signs	Earliest seen&Classif	Algo Based Alternative Syndromes by Probability with Polydactyly at 10 wks, After Exclusion of Meckel-Gruber Syndrome
Polydactyly (101)	10wks	Meckel Gruber Syndrome			
			Sloping forehead		Bandel-Bledj syndrome
			Micrognathia	12wks I	Ellis-Van Creveld syndrome
			Potter-like facies		Short rib polydactyly syndrome Type 1
			Low-set ears	11wks I	Pellier-Killian Syndrome
			Microphthalmia	12wks I	46, XY, -3, +der(5)
			Hypotelorism	10wks I	APERT SYNDROME
			Hypertelorism	15wks	BECKWITH-WIEDEMANN SYNDROME; BWS
			Cleft palate	11wks I	HOLT-ORAM SYNDROME; HOS
			Cleft lip	11wks I	ACHONDROGENESIS, TYPE II; ACG2
			Macrostomia		FRASER SYNDROME
			Short neck		RUTLEDGE LETHAL MULTIPLE CONGENITAL ANOMALY SYNDROME
			Increased NT	10 wks II	SMITH-LEMLI-OPTIZ SYNDROME; SLOS
			Septal defects	11wks I	SIMPSON-GOLAB-BEHMEI SYNDROME, TYPE 1; SGBS1
			Coarctation of aorta	11wks I	ACROCALLOSAL SYNDROME; ACLS
			Pulmonary hypoplasia	15wks	ASPHYXIATING THORACIC DYSTROPHY; ATD
			Splenomegaly		JOUBERT SYNDROME 1; JBTS1
			Asplenia		HYDROLETHALUS SYNDROME
			Accessory spleen		SPLIT-HAND/FOOT MALFORMATION 3; SHFM3
			Single umbilical artery	10wks I	ECTODERMAL DYSPLASIA SYNDROME WITH DISTINCTIVE FACIAL APPEARANCE AND PREAXIAL POLYDACTYLY OF FEET
			Omphalocele	10wks III	VATER ASSOCIATION
			Intestinal malrotation		DANDY-WALKER MALFORMATION WITH POSTAXIAL POLYDACTYLY
					JOUBERT SYNDROME WITH BILATERAL CHORIORETINAL COLOBOMA

Fig. 2C

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[illegible]

**Fig. 2D**

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Teratogens Chart Data Screen

aisaisaisaisaiskajkskaikrja  
super2

sub21  
dr211  
sub22  
dr221  
sub13  
drug131  
super22  
sfdssfs  
sub21  
sub22  
sub13  
super22  
sub21  
sub22  
sub13  
super22  
sub21  
sub22

Week 5 Day 3 Enter

☒ Non-relevant  
☐ Relevant

Save

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30

Fig. 3A

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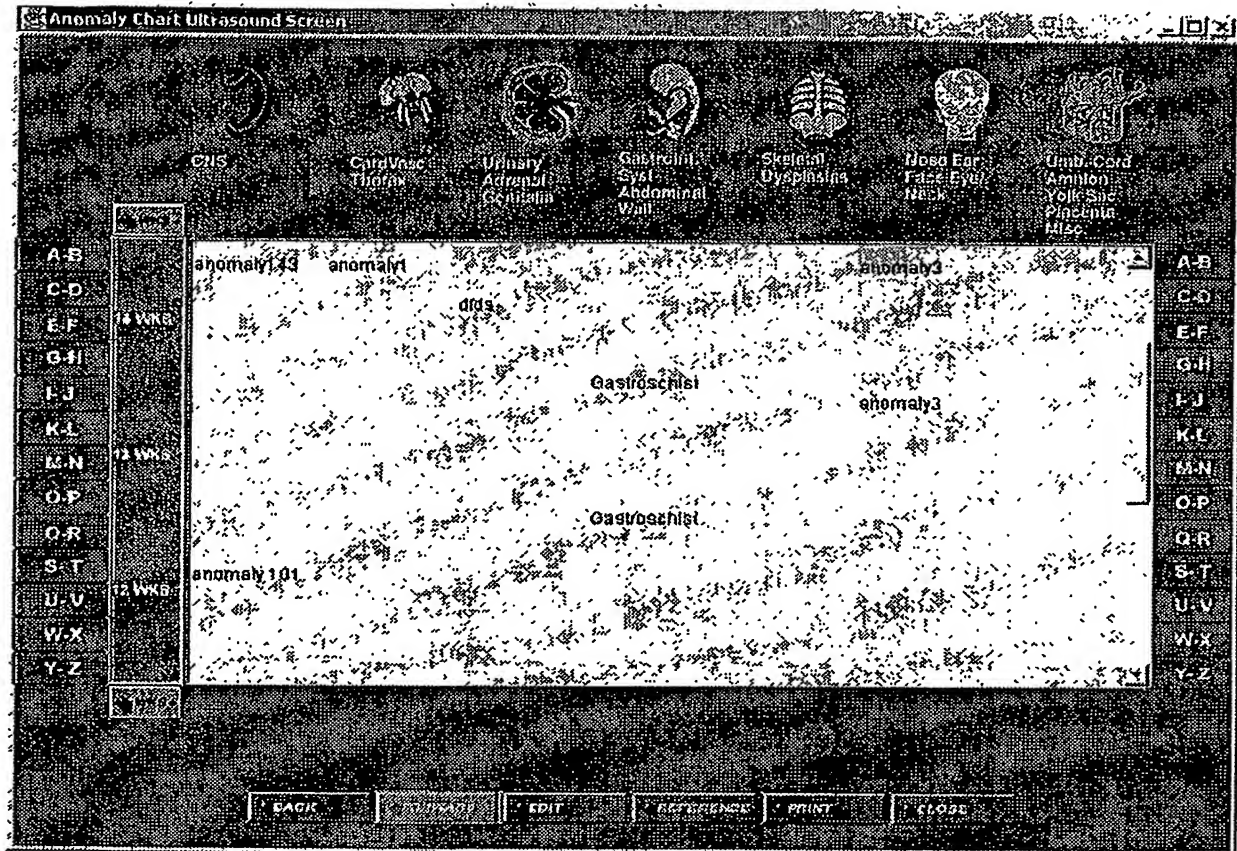


Fig. 3B



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Maternal Disease Chart Screen

42

disease1  
sub11  
sub22  
sub33  
disease2  
sub2.1  
sub2.3  
disease3  
sub3.1  
sub3.2  
sub3.3  
disease4  
sub4.1  
sub4.2  
Disease5  
disease5  
disease5.1  
disease5.2  
Disease6  
dis66

A-B  
C-D  
E-F  
G-H  
I-J  
K-L  
M-N  
O-P  
Q-R  
S-T  
U-V  
W-X  
Y-Z

Week 12 Day 5 Antibodies 0.56 Enter

☐ Low  
☐ High

A-B  
C-D  
E-F  
G-H  
I-J  
K-L  
M-N  
O-P  
Q-R  
S-T  
U-V  
W-X  
Y-Z

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Fig. 4A

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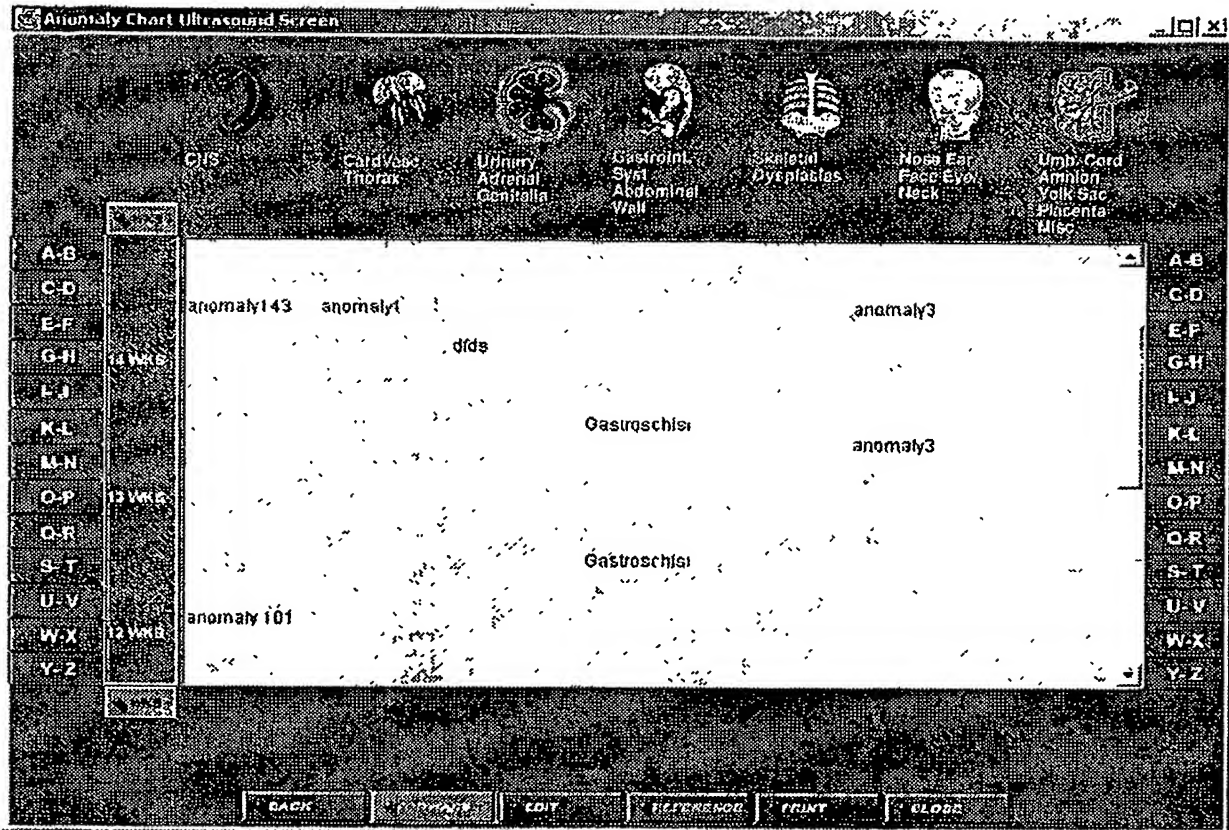


Fig. 4B

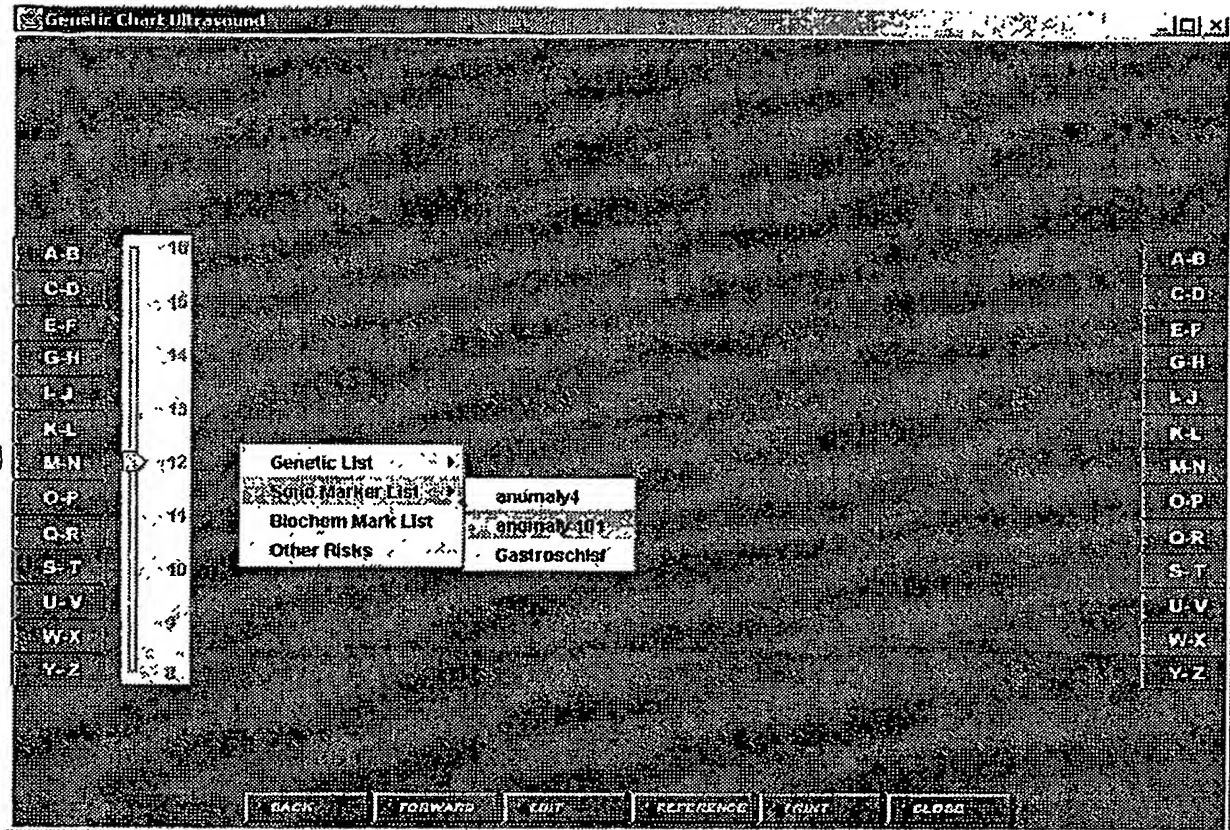


Fig. 4C